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# UNIVERSITÀ DEGLI STUDI DI TORINO

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**Fluorescent in situ hybridization mapping of three fecundity genes on cattle, river buffalo,  
sheep and goat**

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1   **ABSTRACT**

2   One of the goals of molecular cytogenetics applied to livestock is the extension of their genetic  
3   physical maps, especially of loci containing genes related to productions. In this study, a  
4   comparative fluorescence in situ (FISH) mapping of three genes related to fecundity of cattle, river  
5   buffalo, sheep and goat is reported using bovine BAC-clones taking in account the data available on  
6   the BovMap database and considering their physical position and the data obtained from banding  
7   experiments. The following three gene sequences were mapped: tumor necrosis factor- $\alpha$  (TNF),  
8   correlated to male fertility; signal transducer and activator of transcription 5A (STAT5A), important  
9   for its influence on milk production and reproduction activity; melatonin receptor 1A (MTNR1A)  
10   important for reproductive seasonality. BAC probes containing these gene sequences were assigned  
11   by FISH, for the first time, on RB-banded chromosomes of these four important bovids. TNF was  
12   assigned to BTA/CHI23q21-22, OAR20q21-22 and BBU 2p21-22; STAT5A was assigned to  
13   BTA/CHI19q17-21, OAR11q17-21 and BBU3p15-21; MTNR1A was assigned to BTA/CHI27q14-  
14   15, OAR26q14-15 and BBU1p21-22. The three loci were located in homoeologous chromosomes  
15   and chromosome bands, underling the high degree of chromosome homologies among Bovids and  
16   extending the cytogenetic maps of this economically important species.

17

18   **Keywords:** FISH-mapping, fecundity genes, cattle, sheep, goat, river buffalo

19

## 1 INTRODUCTION

2 Cytogenetic mapping is a method used to construct physical maps which are useful tools for  
3 various applications, especially in animal cytogenetics. Indeed, it allows: (a) a precise physical  
4 position on single chromosome bands of both type I and type II loci, especially in bovids using  
5 FISH mapping on R-banded chromosome preparations (Di Meo et al. 2007); (b) to confirm  
6 chromosomes and chromosome regions involved in chromosome abnormalities by using specific  
7 molecular markers (Di Meo et al. 2000; Perucatti et al. 2011; Iannuzzi et al. 2013); (c) to study  
8 chromosome aneuploidies in sperms and oocytes (Pauciullo et al. 2011, 2012; Hornak et al. 2011);  
9 (d) to precisely anchor linkage and RH maps (Stafuzza et al. 2013), as well as genome sequence  
10 contigs to specific chromosome regions (Goldammer et al. 2009).

11 Cattle (*Bos taurus*,  $2n = 60$ , BTA), river buffalo (*Bubalus bubalis*,  $2n = 50$ , BBU), sheep  
12 (*Ovis aries*,  $2n = 54$ , OAR) and goats (*Capra hircus*,  $2n = 60$ , CHI) are very related species from the  
13 evolutionary point of view and, also, the four major domestic bovid species of great economic  
14 importance. Although the location of a lot of genes in these species was identified by linkage and  
15 RH mapping, a small percentage of those loci were physically assigned to the corresponding bands  
16 of specific chromosomal location (Iannuzzi et al. 2003a). So far, several studies on the physical  
17 gene mapping using FISH methodology were reported for cattle, river buffalo, sheep, goat and other  
18 farm animals (Iannuzzi et al. 2003a, 2003b; Di Meo et al. 2007; Schibler et al. 2009).

19 In the present study, three important fecundity genes (TNF, STAT5A and MTNR1A) were  
20 comparatively FISH mapped on cattle, sheep, goat and river buffalo R-banded chromosomes for  
21 first time extending the cytogenetic maps of these species. Tumor necrosis factor- $\alpha$  (TNF) is  
22 correlated to male fertility (Eggert-Kruse et al. 2007; Kocak et al. 2002); signal transducer and  
23 activator of transcription 5A (STAT5A) is important for its influence on milk production and  
24 reproduction activity (Yang et al. 2000; Homer et al. 2013); melatonin receptor 1A (MTNR1A) is  
25 important for reproductive seasonality (Chu et al. 2007; Luridiana et al. 2012).

26

## 1 MATERIALS AND METHODS

2 Peripheral blood samples from cattle (Agerolese breed), sheep (Laticauda breed), goat  
3 (Cilentana breed) and riverbuffalo were cultured and treated for late BrdU and Hoechst 33258  
4 incorporation according to Iannuzzi and Di Berardino (2008). The bovine BAC clones overlapping  
5 studied genes (Table 1) were screened by database searching and ordered from INRA bovine BAC  
6 library (CRB- Biological Resources Centre dedicated to livestock genomics –INRA, Jouy-en Josas,  
7 France) (<http://locus.jouy.inra.fr/cgibin/bovmap/intro2.pl>). Extraction of DNA was done using  
8 CHORI (Children’s Hospital Oakland Research Institute) recommended protocol. DNA was labeled  
9 with biotin and digoxigenin using nick-translation kit (Roche applied science Inc.). Slides were then  
10 treated for FISH with BAC clones overnight in presence of bovine COT-I DNA and sonicated  
11 salmon sperm allocated in a moist chamber. After detection steps with FITC-avidin and anti-  
12 digoxigenin antibodies, Chromosomes were counterstained with Vectashield DAPI H1500 in  
13 Vectashield H 1000 (Vector Lab) antifade solution. Both RB-banding (R-banding by late  
14 incorporation of BrdU) metaphases and fluorescence FITC and TRIC signals were separately  
15 captured by a CCD-camera (Photometrics, cool SNAP, Nikon) and processed by superimposing  
16 FITC and TRIC signals on RB-banding preparations. Chromosome identification and banding  
17 followed the standard karyotypes for cattle, sheep and goat (ISCNDB2000 2001) and river buffalo  
18 (CSKBB 1994).

19

## 20 RESULTS AND DISCUSSION

21 Three major fecundity genes (TNF, STAT5A and MTNR1A), were comparatively  
22 physically FISH-mapped on cattle, sheep goat and river buffalo R-banded metaphase chromosomes  
23 (Figure 1). Loci FISH-mapped with locus name, symbol, clone identification and chromosome  
24 localization are reported in Table 1. TNF maps on BTA/CHI23q21-22, OAR20q21-22 and BBU  
25 2p21-22; STAT5A maps on BTA/CHI19q17-21, OAR11q17-21 and BBU3p15-21; MTNR1A maps  
26 to BTA/CHI27q14-15, OAR11q17-21 and BBU1p21-22. The three loci were located in

homoeologous chromosomes and chromosome bands of the four species extending the cytogenetic maps in these three species chromosomes. FISH-mapping of STAT5A agrees with previous localizations performed in BTA/CHI19 and OAR11 by sequential GTG-banding and FISH (Goldammer et al. 1997), while MTNR1A, earlier assigned to BBU1 by RH-mapping (Miziara et al. 2007) was now assigned to specific chromosome arms and bands (1p21-22).

During the last fifteen years, FISH techniques have been used in domestic animals research mainly to identify chromosomal rearrangements, gene mapping, comparative mapping, and evolutionary chromosome studies. The localization of TNF, STAT5A and MTNR1A on homologous chromosomes and chromosome bands in cattle, sheep, goat and river buffalo (Figure 1; Table 1) confirmed the high conservation of autosomal chromosomes among the bovid species and extended the cytogenetic maps of the four economically important domestic species. However, some discrepancies may exist between the localization of loci reported in a reference genome and the localization obtained by FISH physical mapping, as supported by different papers (De Lorenzi et al. 2010; Partipilo et al. 2011; De Lorenzi et al. 2013). These results clearly indicate the idea that physical localization of genomic elements by FISH can further improve the excellent results obtained by genome sequence projects.

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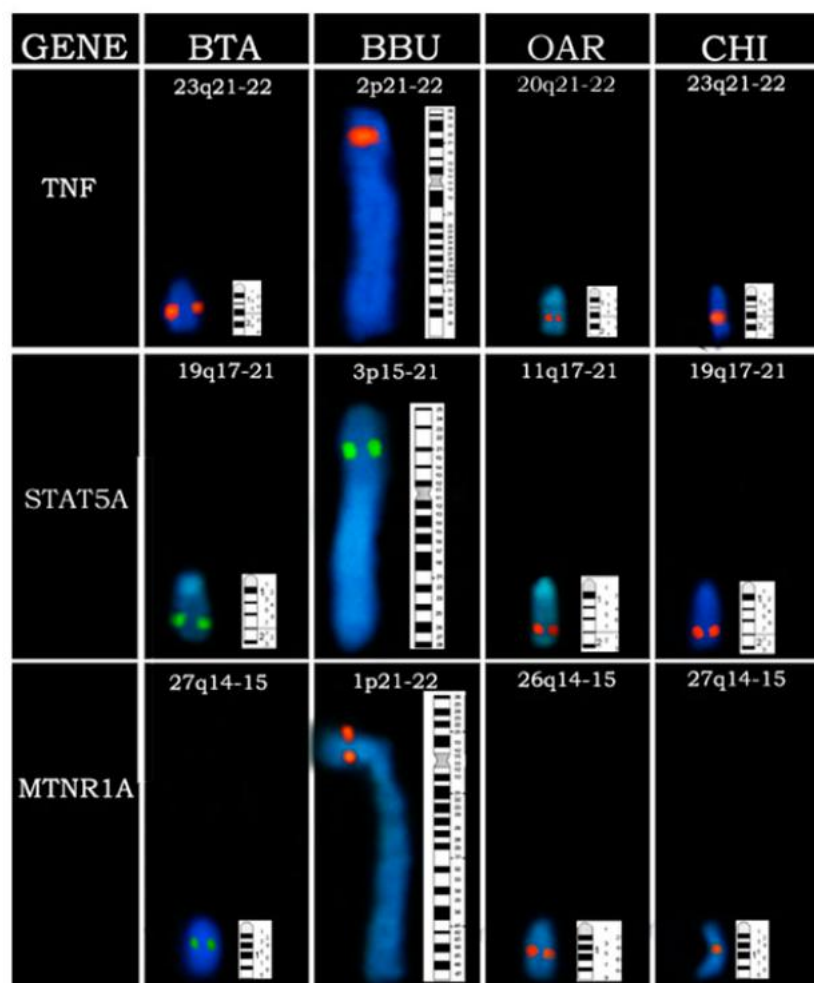


Figure 1. (Color online) Representative FISH results on cattle (BTA), river buffalo (BBU), sheep (OAR) and goat (CHI) chromosomes, using bovine BAC clones containing genes related to fecundity (TNF, STAT5A, MTNR1A). FITC and TRIC signals were superimposed on R-banding chromosomes counterstained with DAPI. For each chromosome, the corresponding standard ideogram (ISCNDB2000 2001; CSKBB 1994) it is also reported.

1 Table 1. BAC-probes, identified DNA sequences of FISH-mapped genes in cattle (BTA), river buffalo (BBU), sheep (OAR), goat (CHI)  
2 chromosomes (ISCNDB2000 2001 and CSKBB 1994), comparison with human (HSA) chromosomes (HGNC).

3

| BAC<br>FISH<br>Probe | Identified DNA sequence<br>within BAC and locus<br>symbol (HGNC) | Gene name  | Cytogenetic localization on RBPI-bands |         |          |          |         |
|----------------------|--|--|--|---------|----------|----------|---------|
|                      |  |  | BTA                                    | BBU     | OAR      | CHI      | HSA     |
| BtINRA-81C03         | <i>TNF</i>   | <i>tumor necrosis factor-<math>\alpha</math></i>               | 23q21-22                               | 2p21-22 | 20q21-22 | 23q21-22 | 6p21.3  |
| BtINRA-243A01        | <i>STAT5A</i>  | <i>signal transducer and activator of<br/>transcription 5A</i> | 19q17-21                               | 3p15-21 | 11q17-21 | 19q17-21 | 17q11.2 |
| BtINRA-448A07        | <i>MTNR1A</i>  | <i>melatonin receptor 1 A</i>                                  | 27q14-15                               | 1p21-22 | 26q14-15 | 27q14-15 | 4q35    |

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